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Journal Title: JOURNAL OF NEUROSCIENCE METHODS
Volume: Volume 341, Number
Publisher: ELSEVIER | 2020-07-15, Pages 108756-108756
Type of Work: Article
Publisher DOI: 10.1016/j.jneumeth.2020.108756
Permanent URL: https://pid.emory.edu/ark:/25593/vtd5f

Final published version: http://dx.doi.org/10.1016/j.jneumeth.2020.108756
Accessed December 2, 2022 10:20 AM EST
Functional network connectivity (FNC)-based generative adversarial network (GAN) and its applications in classification of mental disorders

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Abstract

As a popular deep learning method, generative adversarial networks (GAN) have achieved outstanding performance in multiple classifications and segmentation tasks. However, the application of GANs to fMRI data is relatively rare. In this work, we proposed a functional network connectivity (FNC) based GAN for classifying psychotic disorders from healthy controls (HCs), in which FNC matrices were calculated by correlation of time courses derived from non-
artefactual fMRI independent components (ICs). The proposed GAN model consisted of one discriminator (real FNCs) and one generator (fake FNCs), each has four fully-connected layers. The generator was trained to match the discriminator in the intermediate layers while simultaneously a new objective loss was determined for the generator to improve the whole classification performance. In a case for classifying 269 major depressive disorder (MDD) patients from 286 HCs, an average accuracy of 70.1% was achieved in 10-fold cross-validation, with at least 6% higher compared to the other 6 popular classification approaches (54.5–64.2%). In another application to discriminating 558 schizophrenia patients from 542 HCs from 7 sites, the proposed GAN model achieved 80.7% accuracy in leave-one-site-out prediction, outperforming support vector machine (SVM) and deep neural net (DNN) by 3%–6%. More importantly, we are able to identify the most contributing FNC nodes and edges with the strategy of leave-one-FNC-out recursively. To the best of our knowledge, this is the first attempt to apply the GAN model on the FNC-based classification of mental disorders. Such a framework promises wide utility and great potential in neuroimaging biomarker identification.

Keywords
Resting-state fMRI; Generative adversarial networks (GAN); Deep learning; Classification; Major depressive disorders; Schizophrenia

1. Introduction

Mental disorders cause high socioeconomic burdens and many exhibit comorbidity between each other (Kessler et al., 2012). Machine learning methods based on neuroimaging data have been widely applied in the classification of mental disorder in order to facilitate the diagnosis objectively e.g., support vector machine (SVM), linear discriminant analysis (LDA), and nearest neighbors (NN) (Gao et al., 2018; Jie et al., 2018; Sato et al., 2015; Zhang et al., 2015). In most cases, the raw fMRI data needs to be extracted and processed in order to acquire less redundant and more informative features. Recently, deep learning has achieved remarkable performance in many research field (Arbabshirani et al., 2017; Calhoun and Sui, 2016; LeCun et al., 2015; Yan et al., 2019a; Yang et al., 2019) due to its ability to automatically learns the patterns from the big data without feature selection. For instance, (Kim et al., 2016) adopted a stacked autoencoder to initialize its own weight based on pre-training to increase schizophrenia classification accuracy. (Zeng et al., 2018) investigated the discriminant autoencoder network for multi-site classification of schizophrenia with fMRI. (Yan et al., 2019a) proposed a multi-scale RNN model to enable schizophrenia classification by using time courses of fMRI independent components (ICs) directly. Basically, deep learning methods have demonstrated powerful discriminating ability for classifying mental disorders using fMRI data.

In particular, generative adversarial networks (GAN) have drawn increasing attention due to their capability to perform data generation and have been widely used in the fields of image synthesis, reconstruction, segmentation, and classification (Yi et al., 2019). It has been recently proven successful on several standard classification benchmark tasks (Feng et al., 2019; Salimans et al., 2016; Springenberg, 2015; Vandenhende et al., 2019) with the fact
that the generated samples promote the GAN’s classification ability through adversarial learning, especially in the case of small samples size. However, the high-dimensional data in medical imaging is often with only a limited sample size, which makes it challenging for the classifiers to learn a good decision boundary. In order to improve GAN’s classification performance, feature matching has been implemented as state-of-the-art approaches in many GAN models. Specifically, here feature matching is setting a new objective loss for the generator which is trained to match the feature value in the intermediate layers of the discriminator. It can generate fake samples within the high-density region in feature space, which can split the bounds of different classes because of its continuity to further enhance classification performance (Dai et al., 2017; Salimans et al., 2016).

Till now, the application of GAN on fMRI data is relatively rare. Inspired by this, in this work, we proposed a functional network connectivity (FNC)-based GAN with feature matching for classifying mental disorders from healthy controls (HCs), in which FNC matrices were calculated by correlation of time courses derived from non-artefactual fMRI independent components (ICs). The reason we used FNC is that it is able to reflect functional interactions among structurally segregated brain regions, known as intrinsic functional connectivity networks (ICN) (Calhoun et al., 2014) and reduce the feature dimension greatly. Furthermore, the FNC estimated by independent component analysis (group-ICA) also demonstrated to be more reliable and sensitive in biomarker detection for psychosis (Du et al., 2015; Sui et al., 2018) and can be applied to new individuals (Du et al., 2018). To the best of our knowledge, this is the first attempt to apply GAN on FNC features to discriminate psychiatric patients from controls. We tried on two kinds of disorders: 1) MDD vs HC (555 subjects, 269 MDD, 286 HCs); 2) SZ vs HC (1100 subjects, 558 SZ, 542 HCs). The proposed GAN model combines fake FNC generation and classification together into a unified optimization framework. To increase result interpretability, leave-one-FNC-out looping was adopted to identify the most contributing FNCs.

2. Materials and methods

2.1. Generative adversarial network with feature matching

The proposed GAN model consisted of one discriminator (real FNCs) and one generator (fake FNCs), each has four fully-connected layers. Furthermore, feature matching, the generator was trained to match the feature value in intermediate layers of the discriminator, was implemented by specifying a new objective loss for the generator to improve classification performance. FNC estimated for each subject based on group-ICA was used as the input of the GAN module. To verify that adversarial training (AD) and feature matching (FM) in GAN do directly improve the learning capacity of the GAN method, the ablation experiments were performed in the paper. Equations are listed in the Supplementary files for a precise definition of the GAN model with feature matching.

2.1.1. Generative adversarial network—GAN may learn a better decision boundary than the traditional methods because the generated samples promote the discriminator’s classification ability through adversarial learning, especially in the case of a small sample
As shown in Fig. 1, a GAN architecture was applied for classification. The GAN model was composed of a discriminator and a generator with four fully-connected layers, optimizing its network parameters through continuous competitions. The discriminator’s output layer has K+1 classes, where K = 2 for the real class from data x and K-1+ class for the generated image. It is noteworthy to note that the discriminator can realize multiclass discriminative ability by a soft-max classifier. Compared to the unsupervised GAN model, the proposed GAN incorporated both labeled and generated data into the loss function.

### 2.1.2 Feature matching

Feature matching was able to address the instability of GAN by specifying a new objective loss for the generator, which can prevent it from overtraining on the discriminator (Salimans et al., 2016). Instead of directly maximizing the output of the discriminator, the new objective enabled the generator to generate data that matched the statistics of the real data, where we only used the discriminator to specify the statistics that were worth matching. The generator was trained to match the feature value in intermediate layers of the discriminator. The intermediate layer we set in this study is the third hidden layer. Through feature matching by the generator, discriminative features can be found to distinguish real data from generated data by training the discriminator. It can generate fake samples within the high-density region in feature space, which can split the bounds of different classes because of its continuity to further enhance classification performance (Dai et al., 2017).

Activations in an intermediate layer of the discriminator were denoted as \( f(x) \), our new objective for the generator can be defined as follows:

\[
\| E_x \sim p_{\text{data}} f(x) - E_z \sim p_z(G(z)) \|_2^2
\]

Where \( x \) represents the input data, \( p_{\text{data}} \) and \( p_z \) represent the distribution of the input data and the distribution of the input noise, \( G \) is the generator, and \( D \) is the discriminator.

### 2.2 GAN model implementation

The GAN model was trained and evaluated by using Theano and cikit-learn (https://scikit-learn.org/). The GAN model consisted of one discriminator and one generator that had four layers respectively. The output layer of the discriminator has \( K+1 \) classes, where \( K = 2 \) for the input pattern from HC and patient group, and the \( [K+1]_{th} \) class is for generating images. The high-dimensional features of each subject may lead to overfitting on the training set. In order to reduce the overfitting susceptibility, L2 normalization (\( L2 = 0.1 \)) and batch normalization were added to the generator to improve modal generalization. In addition, weight normalization (weight normalization = 0.1) was used to the output of each layer of the discriminator to prevent overfitting.

The Adam optimizer was adopted as minimizing the loss of the GAN model and a standard error back-propagation algorithm was used by training the GAN model with multiple layers. The batch size was set as 120 in the training process. In addition, to overcome overfitting, the weights were controlled with weight norm regularization. Different layers were attempted to the constructed architecture of the GAN model and results revealed that using four layers could obtain the optimal classification performance. The learning rate was set as
In this experiment, the training time for GAN was about one hour, while the testing time for a new subject is less than 1 s. All trainings and experiments were completed on a standard workstation (Intel(R) Xeon(R) CPU E5–1650 v4 @ 3.60 GHz, 6 CPU cores, 12GB NVIDIA GTX TITAN).

2.3. **GAN classification framework**

In this study, k-fold multi-site pooling classification and leave-one-site-out classification were conducted. In order to verify the validity of the proposed GAN method, five conventional methods SVM, NN, Gaussian Process, Naive Bayes, and a deep learning method DNN were used as a comparison in this study.

10-fold cross-validation was applied to verify the generalization ability of the classifiers in the multi-site pooling classification. The training dataset and the testing dataset were embedded in nested 10-fold cross-validation cycles. As for all the models, we used nine folds as the training set, and one fold for the testing dataset. A specific imaging site was applied as the testing set and the sample of other sites as the training set in the leave-one-site-out transfer classification. Accuracy (ACC), sensitivity (SEN), specificity (SPE), F-score (F1) and area under curve (AUC) verify the classification performance. The formula is defined as follows:

\[
\text{ACC} = \frac{TP + TN}{TP + TN + FP + FN}, \quad \text{SPE} = \frac{TN}{TN + FP}, \quad \text{SEN} = \frac{TP}{TP + FN}, \quad \text{PPV} = \frac{TP}{TP + FP}, \quad \text{F1} = \frac{2 \cdot \text{SEN} \cdot \text{PPV}}{\text{SEN} + \text{PPV}}
\]

Where true positive, true negative, false positive, false negative and positive predictive value was denoted as TP, TN, FP, FN, PPV respectively. The means and standard deviations of ACC, SEN, SPE, F1 and AUC were obtained by 10-time 10-fold cross-validation and a two-sample t-test was adopted as the comparison of the performance of different classification models.

2.4. **Estimate the contributing FNC (Leave-one-FNC-out)**

The basic idea is that potential biomarkers are such that the removal of these features leads to the most significant degradation of accuracy. The specific implementation is as follows:

In order to calculate the most contributing FNC, we substitute the $d_h$ FNC with default value 0 while keeping other FNC as they were. This is equivalent to erasing the contribution of $d_h$ FNC. The classification accuracy with reduced FNC may decrease compared to that using all FNCs. Compared to that using all features, the classification accuracy of the trained model which is fed with reduced features may decrease. Based on the leave-one-feature-out strategy, we sorted the classification accuracy when getting rid of each FNC. Then the FNCs which cause the top 3% decrease of classification accuracy were recorded and considered as the most contributing features. Similarly, for the most contributing FNC node, we summarized the top 30% contributing FNCs and discovered the top 10% nodes that linking most of the FNCs (higher degrees). Moreover, the selected features are visualized with

The brain regions in MDD were grouped into seven functional subsystems for visualization based on their anatomical and functional properties: auditory (AUD), cognitive control (CC), sensorimotor network (SMN), default-mode network (DMN), frontal network (FN), visual (VIS), and cerebellum (CB).

The brain regions in SZ were grouped into eight functional subsystems for visualization based on their anatomical and functional properties: sub-cortical (SC), auditory (AUD), Sensorimotor network (SMN), visual (VIS), default-mode network (DMN), attentional network (ATN), frontal network (FN), and cerebellum (CB). long term, we should look at both analyzed via neuromark so we can directly compare different mental disorders such as MDD and SZ.

3. Participants

In this study, 555 Chinese Han subjects including 269 MDD patients and 286 HCs derived from 4 sites were used, including the Henan Mental Hospital of Xinxiang (Site 1), the West China Hospital of Sichuan (Site 2), the Anding Hospital of Beijing (Site 3), and the First Affiliated Hospital of Zhejiang (Site 4). No significant group difference between HC and MDD was obtained in age or gender (age: p = 0.22; gender: p = 0.18). The DSM-IV based on SCID-P interviews was used to diagnose patients. HCs were interviewed using SCID-I/NP, and the first-degree relatives with any mental illness were excluded. Each site acquired approval from their respective research ethics boards, and informed consent was received from each subject prior to scanning by each site’s Institutional Review Boards. Patients were excluded if they had a history of head injury which cause significant non-psychiatric medical illness, or unconsciousness for longer than a few seconds. These individuals were also excluded with obsessive-compulsive disorder, posttraumatic stress disorder or an active substance use disorder. Table 1 provides a statistical information, please see more information in Table S1.

For SZ data, 1100 Chinese subjects (542 HCs and 558 SZ patients) from seven sites were used, including the Peking University sixth Hospital (Site 1), the Huilongguan Hospital (Site 2), the Henan Mental Hospital (Site 3), the Henan Mental Hospital (Site 4), the Xijing Hospital (Site 5), the Renmin Hospital of Wuhan University (Site 6), and the Zhumadian Psychiatric Hospital (Site 7). Each site received approval from the relevant Ethics Committees and all study participants obtained written informed consents. All the SZ patients were evaluated based on the SCID and diagnosed by experienced psychiatrists by using DSM-IV-TR. All the HCs which were free of Axis I or II disorders were recruited from the same local geographical areas as the patients cohort through local advertisement. The exclusion criteria were suitable for all subjects included substance abuse or dependence, pregnancy, prior electroconvulsive therapy, and current or past neurological illness. Table 2 provides a statistical information, please see more information in Table S2.
3.1. **Data acquisition**

The resting-state fMRI data for MDD patients and demographically matched HCs were collected on a 3 T Philips scanner at Site 2 and on a 3 T Siemens scanner (Verio, Germany) at site 1, site 3 and site 4. The field of view (FOV) was 240 × 240 mm (64 × 64 matrix) at site 2, 220 × 220 mm (64 × 64 matrix) at site 1 and site 4, 200 × 200 mm (64 × 64 matrix) at site 3 and all sites have 240 volumes of echo planar images obtained by the same parameters: repetition time (TR) = 2000 ms; echo time (TE) = 30 ms; flip angle (FA) = 90. Site 1 was 33 sequential ascending axial slices of 4 mm thickness, Site 3 was 33 sequential ascending axial slices of 3.5 mm thickness and site 2, and site 4 were 38 sequential ascending axial slices of 4 mm thickness. During scanning, head movement and scanner noise were minimized by foam padding and earplugs and subjects followed the instructions to keep eyes closed and stay awake when lying still.

The resting-state fMRI data for SZ patients and age- and gendermatched HCs were collected on a 3.0 T Siemens Trio Tim Scanner in Site 1, site 2 and site 5, a 3.0 T Siemens Verio Scanner at site 3, and a 3.0 T Signa HDx GE Scanner at site 4, site 6, and site 7. During scanning, head movement and scanner noise were minimized by foam padding and earplugs and subjects followed the instructions to keep eyes closed and stay awake while lying still.

3.2. **Data preprocessing and FNC measure**

The SPM8 (http://www.fil.ion.ucl.ac.uk/spm/) software is used by the preprocessing of the resting-state fMRI data and the preprocessing flow is as follows: The first 10 volumes were discarded to exclude T1 equilibration effects, slice timing correction, motion correction, normalization in the standard Montreal Neurological Institute (MNI) space (3 mm isotropic voxels), denoising and spatially smoothing with an 8 mm full-width half max Gaussian kernel. Each voxel time course was zscored to normalize variance across space. In addition, to limit the impact of head motion, a maximum translation of > 2 mm or rotation of > 2 mm or framewise displacements (FD) > 1 mm was excluded in the subjects. Results indicate that mean FD for all subjects were < 0.5 mm and there was no significant group difference between HC vs SZ patients and a significant group difference between HC vs MDD patients on mean FD (HC: 0.14 ± 0.070, MDD: 0.12 ± 0.066, two-sample t-test: p = 0.001, HC: 0.137 ± 0.071, SZ: 0.142 ± 0.085, two-sample t-test: p = 0.98). For MDD and HC, since the mean FD showed a significant group difference, we further regressed out the mean FD from each functional network connectivity to eliminate the potential group difference (Yuan et al., 2016). Moreover, we regressed age, gender in order to eliminate the effects of these variables.

For MDD patients and HCs, the fMRI data were decomposed into subject-specific spatial ICs and its corresponding time courses using a spatially constrained ICA back-reconstruction approach called group ICA implemented in the GIFT software (http://trendscenter.org/software/gift) which is robust to artifacts (Du et al., 2016), resulting in 29 selected ICNs from 100 group independent components. More details of spatial maps are listed in the Supplementary Figure. S3. ICA was applied first to decompose the 4D fMRI into useful ICNs and some noise ICs that were thrown off, which is also a denoising step. Due to the time courses of the selected ICNs may still contain remaining noise sources, the
additional post-processing including detrending, controlling covariates by regression, and bandpass filtering was applied (Allen et al., 2012). The time courses of selected ICNs were post-processed by detrending linear, quadratic and cubic trends, regressing out 6 realignment parameters and their temporal derivatives, despiking, and bandpass filtering between [0.01~0.15] Hz using a 5th order Butterworth filter. FNC was computed as the pairwise correlation between any two ICN time courses for each subject, which was further used as input feature of the GAN model.

For SZ patients and HCs, the fMRI data were decomposed into subject-specific spatial ICs and its time courses by performing group-ICA within the GIFT software (Du et al., 2016), resulting in 50 selected ICNs from 100 group independent components. More details of spatial maps are listed in the Supplementary Figure. S4. The time courses (TCs) of selected ICNs are post-processed by detrending, regressing out head motion, despiking and lowpass filtering (<0.15 Hz) and the FNC matrices are calculated as the Pearson’s correlation in each pair of ICs, which was further used as input feature of the GAN model.

4. Results

4.1. Ten-fold and leave-one-site-out classification in MDD/HC

To demonstrate the performance of the proposed method, we compared the proposed method with other state-of-art methods, including five conventional methods SVM, NN, Gaussian Process, Naive Bayes, and AdaBoost and a deep learning method DNN. The training dataset and the testing dataset were embedded in nested 10-fold cross-validation cycles. As for all the models, we used nine folds as the training set, and one fold for the testing dataset. In order to test the generalization of the model, we train the different models with the leave-one-site-out method. The means and standard deviations of ACC, SEN, SPE, F1 and AUC were obtained by 10-time 10-fold cross-validation and a two-sample t-test was adopted as the comparison of the performance of different classification models.

The classification results of the GAN model compared with the other six methods were summarized in Table 3, Table 4, and Table S3. The accuracy of 70.1 ± 0.6% and 64.3 ± 2.9% was obtained by using the GAN model in the 10-fold cross-validation and leave-one-site-out method, which is significantly higher than those obtained by using those traditional classification methods and DNN approach. For the binary classification problem, the results suggested that the proposed GAN method significantly outperformed five conventional methods SVM, NN, Gaussian Process, Naive Bayes, and AdaBoost and a deep learning method DNN in terms of ACC, SEN, SPE, F1, and AUC measures (P < 0.05, two-sample t-test), and achieved good generalization results in the leave-one-site-out method (Table 3, Table 4, Table S3, Fig. 2A and B) on account of our GAN model makes full use of adversarial learning and feature matching that alleviate the small size problem of FNC images and mitigate the instability of GAN. The t-distributed stochastic neighbor embedding (t-SNE) was used to visualize the GAN classification performance. The t-SNE visualization result as shown in Fig. S1, indicating that the proposed GAN model shows better separating power and can provide intuition interpretation for the increase of classification performance. In addition, the result that generated samples by the GAN method in comparison with the true FNC as shown in Fig.S2. The analysis of the comparison is listed in the Supplementary
files, revealing that the distribution of FNC generated by GAN is similar to the distribution of real FNC.

4.2. The classification of ten-fold and leave-one-site-out in SZ/HC

To validate the effectiveness of the GAN model, we evaluated the proposed GAN method on the task of SZ classification (542 HCs and 558 SZ). The proposed method was compared with six state-of-art methods using 10-fold cross-validation and leave-one-site-out methods. Results were summarized in Table 5, Table 6, and Table S4, the accuracy of 82.1 ± 0.7% and 80.7 ± 3.8% was obtained by using the GAN model in the 10-fold cross-validation and leave-one-site-out method, demonstrating that the classification performance of GAN was significantly better than other methods (Table 5, Table 6, Table S4, Fig. 2C and Fig. 2D). The proposed GAN model can be generalized to classify the new site. In addition, the t-SNE visualization result is shown in Fig. S1.

4.3. Most contributing FNC for MDD-HC and SZ-HC classification

The research intention of the fMRI classification is that a specific set of features may be used to diagnose the mental disorders as stable biomarkers and to ensure different datasets have a consistent result. In addition, the interpretability of deep learning has always been one of the factors limiting the application of deep learning in the medical field instead of classification performance. Recently, some research has evaluated the interpretability of deep learning (Kim et al., 2016; Yan et al., 2019b; Zeng et al., 2018). The basic idea is that potential biomarkers are such that the removal of these features leads to the most significant degradation of accuracy. Here the most contributing FNC for classification was analyzed in the multi-site pooling strategy with the leave-one-FNC-out method. It was found that the most important FNCs for MDD are presented in Fig S5.a and Table 7. We also evaluated the proposed method on the classification of SZ (542 HCs and 558 SZ) using the leave-one-FNC-out method. It was found that the most important FNCs for MDD are presented in Fig S5.b and Table 8. Moreover, to present the functional “network” connectivity feature, the BRANT (Xu et al., 2018) was applied to plot these brain maps which contain the most contributing FNCs related to a specific network and these brain maps were presented in Fig. 3 and 4.

5. Ablation study

To verify that adversarial training (AD) and feature matching (FM) in GAN do directly improve the learning capacity of the GAN method, we performed the ablation experiments by removing either adversarial training or feature matching, similar as did in (Huang et al., 2016). Specifically, we compared the performance of GAN models with adversarial training vs no adversarial training (No-AD) (Fig. 5a, b) and feature matching vs no feature matching (No-FM) (Fig. 5c,d). We obtained the means and standard deviations of ACC, SEN, SPE, F1 and AUC by 10-time 10-fold cross-validation and a two-sample t-test was adopted as the comparison of the classification performance. It is clear that the GAN model with adversarial training outperformed the GAN with No-AD in almost all evaluation criteria, with a 6% increase in MDD. Moreover, the GAN model with FM also achieved better
classification accuracy compared to the GAN with No-FM (70% vs 69% in MDD, 82 vs.
81% in SZ).

6. Discussion

In this work, we developed a novel GAN model using resting-state fMRI to discriminate
mental disorders from HCs in large, multi-site datasets. Our model alleviates the small size
problem of FNC images by making full use of generated FNC samples. In addition, the
proposed GAN model defines adversarial objections between the generators and
discriminator, which uses adversarial learning and feature matching to further improve the
classification performance of the discriminator. Results showed that the GAN model
achieved an average 70.1% accuracy with 10-fold cross-validation in MDD vs HC, at least
6% higher than five conventional methods and deep neural net (DNN) (54.5–64.2%). To
validate the effectiveness of the GAN model, we further applied it to a large-scale multi-site
schizophrenia (SZ) dataset including 558 patients and 542 HCs from seven sites, achieving
80.7% accuracy in leave-one-site prediction, outperforming SVM and DNN by 3–6%,
demonstrating the efficacy of the GAN approach. Subsequently, to increase result
interpretability, leave-one-FNC-out looping was adopted, and results suggested the most
contribute FNCs were primarily located in the FN, DMN, CB, SMN, and CC for MDD
and in the DMN, SC, VIS, and SMN for SZ. Thus, such a framework promises wide utility
and may have great potential in neuroimaging biomarker identification. In this study the
good classification performance may be attributed to the following aspects:

This is the first attempt to apply GAN on FNC-based classification, results from ablation
experiments suggest that the generated samples in adversarial training did promote the
classification ability for FNC-based GAN, especially in the case of a small sample size. The
potential reason might be that GAN acts as a regularizer for the decision surface of the
classifier to improve the classification performance (Vandenhende et al., 2019). Comparing
with the conventional DNN network without the adversarial item in the cost function, we
found that the accuracy of the GAN is significantly better than the conventional DNN
network with a 6% increase for MDD and HC.

Second, feature matching (FM) in GAN could directly improve the classification
performance of the GAN model. We compared the performance of GAN models with
feature matching vs no feature matching (No-FM). As shown in Fig. 5, the GAN model
showed increased performance from almost all evaluation criteria. In this study, the
leaveone-site-out transfer classification also gained promising accuracies indicating that the
classification models are independent with imaging sites. Maybe due to no prior knowledge
from the testing site, the leaveone-site-out transfer classification accuracies were lower than
the accuracies of multi-site pooling classification to some extent. However, the present
results suggest that the GAN method proposed may provide a potential realistic solution for
individual diagnostic classification with MDD vs HC and SZ vs HC across independent
imaging sites.

As to the most contributing FNC, results showed that the most contributing features were
primarily located in FN, DMN, CB, SMN, and CC in the classification of MDD patients.
involving in cognitive function and emotion regulation (Kaiser et al., 2015; Mulders et al., 2015). The research (Mulders et al., 2015) is consistent with the result of the current study that patients of the MDD observed abnormal FNCs between DMN and CC, CB, which are related to frontal, parietal, and postcentral gyrus. The DMN and cerebellum are related to self-processing and emotional (Drevets et al., 2008; Sheline et al., 2010), and the FN network has an important influence on cognitive ability and mood regulation (Drevets et al., 2008; Steele and Lawrie, 2004). The imbalance within externally-directed attention and control systems was caused by the aberrant connectivity of the MDD between anterior cingulate (AC) and inferior parietal lobule (IPL) (Kaiser et al., 2015) and the perception and emotional facial expressions processing was related to the aberrant FNCs in AUD and FN located in inferior temporal gyrus (ITG), superior frontal gyrus (SFG) and middle frontal gyrus (MFG) (Haxby et al., 2000). Abnormal reactivity caused by FNCs in precuneus and fusiform gyrus (FFG) might accept as a depression early biomarker as described in (Hahn et al., 2011). The SFG is related to executive memory emotions (Fuster, 2001; Jaworska et al., 2015) and the motor, premotor and prefrontal networks are related to executive memory. Executive function abnormalities in MDD patients were implied by the aberrant FNCs with SFG in FN and ITG (Fuster, 2001; Wang et al., 2008).

Results suggested that the most contributing FNCs were primarily located in DMN, SC, VIS, and SMN in the classification of SZ patients involving in deficits in a multitude of cognitive domains (Sheffield and Barch, 2016). Abnormality of the DMN has been linked to SZ in past studies (Sui et al., 2013). The aberrant FNC is associated with angular gyrus (AG) and precuneus in the DMN, which may cause episodic memory deficits and auditory hallucinations in SZ patients (Mondino et al., 2015; Vercammen et al., 2010). Recent study manifest that the deficiency of working memory in SZ is also linked with deficient DMN suppression (Pu et al., 2016). The delusional thought and hallucinations may be caused by precuneus and putamen in the SC network (Braff, 1993; Cui et al., 2016). Hence, SC-related FNC alteration together with DMN abnormality which underlies impaired attention may have an important effect on the positive symptoms of SZ (Camchong et al., 2009; Unschuld et al., 2013). The SMN with postcentral gyrus (PoCG) and precentral gyrus (PreCG) has been shown to be related to higher-order cognitive functions and its connectivity alterations have been associated with cognitive and social-cognitive deficits of SZ in past studies (Hooker et al., 2012; Kaufmann et al., 2015). The FFG is regarded as the critical factor for face recognition, which may possibly be associated with impaired facial recognition in SZ.

Several limitations should be mentioned for this study. One is we used only resting fMRI, however, classifying complex disorders may become more reliable and precise by applying multi-modal neuroimaging fusion (Cetin et al., 2016; Sui et al., 2020, 2011). Second, the generalizability of the classification framework to scanners outside our harmonized study can not be evaluated. Third, it is difficult to assess the medication effect due to the limited sample size and the information inadequacy of antipsychotic or emotion stabilizing medications. Finally, data preprocessing may be vital for machine learning, and each preprocessing step of brain imaging may have a potential effect on the final accuracy in deep learning. Thus, it is important to evaluate the influence of data preprocessing for deep learning in future, and the dynamics of functional connectivity may be another option to improve classification (Meng et al., 2017; Rashid et al., 2016; Zhi et al., 2018).
In summary, to the best of our knowledge, this is the first attempt to apply GAN on FNC-based classification, which integrates both adversarial training and feature matching. Application on discriminations of MDD-HC and SZ-HC verified the effectiveness of the proposed method. Compared with five traditional classification methods and a DNN approach, the GAN model achieved at least 6% higher in MDD classification and 2% higher in SZ classification, suggesting its utility as a potentially powerful tool to aid in discriminative mental disorder detection.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

This work was supported by the Natural Science Foundation of China (No.61773380), the Strategic Priority Research Program of the Chinese Academy of Sciences (No. XDB32040100), Beijing Municipal Science and Technology Commission (Z181100001518005), the National Institute of Health (R01MH117107, R01EB005846, and P20GM103472) and the National Science Foundation (1539067). The authors report no financial interests or potential conflicts of interest.

**References**


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J Neurosci Methods. Author manuscript; available in PMC 2020 July 17.


Fig. 1. Overview of the GAN model for MDD vs HC and SZ vs HC classification. The GAN model was composed of a discriminator and a generator with four fully-connected layers. FNC estimated for each subject based on group-ICA was used as the input of the GAN module.
Fig. 2.
The results of multi-site pooling classification and leave-one-site-out transfer classification for MDD vs HC and SZ vs HC.
(a) The MDD classification results of 10-fold multi-site pooling classification. (b) The MDD classification results of leave-one-site-out transfer classification. (c) The SZ classification results of 10-fold multi-site pooling classification. (d) The SZ classification results of leave-one-site-out transfer classification. In all cases, * means p < 0.05 in two-sample t-tests.
Fig. 3.
Overview of the most contributing FNC related to a specific network for MDD vs HC. AUD: auditory; CC: cognitive control; SMN: sensorimotor network; DMN: default-mode network; FN: frontal network; VIS: visual; CB: cerebellum; ITG: inferior temporal gyrus; MFG: middle frontal gyrus; MTG: middle temporal gyrus; SFG: superior frontal gyrus; AC: anterior cingulate; IPL: inferior parietal lobule; STG: superior temporal gyrus; PoCG: postcentral gyrus; Precun: precuneus; MeFG: medial frontal gyrus; FFG: fusiform gyrus.
Fig. 4.
Overview of the most contributing FNC related to a specific network for SZ vs HC. SC: subcortical; AUD: auditory; SMN: Sensorimotor network; VIS: visual; DMN: default-mode network; ATN: attentional network; FN: frontal network; CB: cerebellum; PoCG: postcentral gyrus; MOG: middle occipital gyrus; AG: angular gyrus; PreCG: precentral gyrus; STG: superior temporal gyrus; FFG: fusiform gyrus.
Fig. 5.
Effect of adversarial training and feature matching.
(a) The comparison of AD vs NO-AD in MDD classification. (b) The comparison of AD vs NO-AD in SZ classification. (c) The comparison of FM vs NO-FM in MDD classification. (d) The comparison of FM vs NO-FM in SZ classification. FNC, Functional network connectivity; AD: adversarial training; FM: feature matching. In all cases, * means p < 0.05 in two-sample t-tests.
### Table 1

Demographic information of the MDD/HC database.

<table>
<thead>
<tr>
<th></th>
<th>MDD</th>
<th>HC</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of scans</td>
<td>269</td>
<td>286</td>
<td>NA</td>
</tr>
<tr>
<td>Age (mean ±std, yrs)</td>
<td>32.8±10.6</td>
<td>31.7±10.4</td>
<td>0.22</td>
</tr>
<tr>
<td>Gender (M/F)</td>
<td>105/164</td>
<td>105/181</td>
<td>0.18</td>
</tr>
<tr>
<td>SYMPTOM SEVERITY HDRS</td>
<td>18.3±8.3</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>BDI</td>
<td>20.4±6.9</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

Table 1 Note: MDD: major depressive disorder; HC: healthy control; yrs: years; BDI: beck depression rating scale; std: standard deviation; HAMD: Hamilton depression scale; F: female; M: male; NA: not applicable.
Table 2

Demographic information of the SZ/HC database.

<table>
<thead>
<tr>
<th></th>
<th>HC</th>
<th>SZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of scans</td>
<td>542</td>
<td>558</td>
</tr>
<tr>
<td>Age (mean ±std, yrs)</td>
<td>28.0 ± 7.2</td>
<td>27.6 ± 7.1</td>
</tr>
<tr>
<td>Gender (M/F)</td>
<td>276/266</td>
<td>292/266</td>
</tr>
</tbody>
</table>

Table 2 Note: SZ: schizophrenia; HC: healthy control; yrs: years; std: standard deviation; F: female; M: male.
### Table 3

Performance of different methods in MDD/HC classification with 10-fold cross-validation.

<table>
<thead>
<tr>
<th>Method</th>
<th>ACC(%)</th>
<th>SEN(%)</th>
<th>SPE(%)</th>
<th>F1(%)</th>
<th>AUC(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nearest Neighbors</td>
<td>54.5(0.9)</td>
<td>55.9(0.9)</td>
<td>53.1(0.9)</td>
<td>56.1(0.9)</td>
<td>54.5(0.9)</td>
</tr>
<tr>
<td>AdaBoost</td>
<td>54.6(1.7)</td>
<td>55.8(1.5)</td>
<td>53.3(1.8)</td>
<td>56.6(1.8)</td>
<td>54.5(1.7)</td>
</tr>
<tr>
<td>Naive Bayes</td>
<td>59.2(0.9)</td>
<td>62.1(1.0)</td>
<td>56.8(0.8)</td>
<td>57.3(1.0)</td>
<td>59.4(0.9)</td>
</tr>
<tr>
<td>Gaussian Process</td>
<td>60.4(0.6)</td>
<td>61.1(0.8)</td>
<td>59.6(0.5)</td>
<td>62.4(0.4)</td>
<td>60.3(0.6)</td>
</tr>
<tr>
<td>Linear SVM</td>
<td>62.8(0.7)</td>
<td>61.9(0.6)</td>
<td>64.2(0.9)</td>
<td>66.7(0.7)</td>
<td>62.5(0.7)</td>
</tr>
<tr>
<td>Deep Neural Net</td>
<td>64.2(0.9)</td>
<td>64.4(1.0)</td>
<td>64.1(1.2)</td>
<td>66.3(1.2)</td>
<td>64.1(0.9)</td>
</tr>
<tr>
<td>GAN</td>
<td>70.1(0.6)</td>
<td>73.5(4.7)</td>
<td>66.5(4.7)</td>
<td>71.7(1.5)</td>
<td>70.3(0.9)</td>
</tr>
</tbody>
</table>

Table 3 The performance of different methods in MDD/HC classification with 10-fold cross-validation. SVM, support vector machine; GAN, Generative Adversarial Networks; ACC, Accuracy; SEN, sensitivity; SPE, specificity; F1, F-score; AUC, area under curve; MDD: major depressive disorder; HC: healthy control.
Table 4

Performance of the leave-one-site-out method in MDD/HC classification.

<table>
<thead>
<tr>
<th>Method</th>
<th>ACC(%)</th>
<th>SEN(%)</th>
<th>SPE(%)</th>
<th>F1(%)</th>
<th>AUC(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nearest Neighbors</td>
<td>55.5(2.0)</td>
<td>57.3(2.5)</td>
<td>53.9(4.6)</td>
<td>55.5(4.4)</td>
<td>55.6(1.2)</td>
</tr>
<tr>
<td>AdaBoost</td>
<td>47.9(1.4)</td>
<td>49.3(5.2)</td>
<td>47.1(1.5)</td>
<td>42.1(1.6)</td>
<td>48.3(2.0)</td>
</tr>
<tr>
<td>Naive Bayes</td>
<td>55.1(6.8)</td>
<td>59.5(9.5)</td>
<td>52.8(6.9)</td>
<td>48.2(12.7)</td>
<td>55.6(6.3)</td>
</tr>
<tr>
<td>Gaussian Process</td>
<td>56.9(5.0)</td>
<td>59.2(4.6)</td>
<td>55.0(6.8)</td>
<td>55.8(8.9)</td>
<td>57.1(2.6)</td>
</tr>
<tr>
<td>Linear SVM</td>
<td>55.9(6.2)</td>
<td>59.0(2.7)</td>
<td>53.7(9.0)</td>
<td>52.4(15.8)</td>
<td>56.1(4.8)</td>
</tr>
<tr>
<td>Deep Neural Net</td>
<td>55.0(3.4)</td>
<td>60.8(5.0)</td>
<td>52.4(3.1)</td>
<td>44.7(2.9)</td>
<td>55.6(1.8)</td>
</tr>
<tr>
<td>GAN</td>
<td>64.3(2.9)</td>
<td>61.5(3.1)</td>
<td>70.8(6.5)</td>
<td>70.5(1.5)</td>
<td>63.8(3.4)</td>
</tr>
</tbody>
</table>

Table 4 The performance of the leave-one-site-out method in MDD/HC classification. SVM, support vector machine; GAN, Generative Adversarial Networks; ACC, Accuracy; SEN, sensitivity; SPE, specificity; F1, F-score; AUC, area under curve; MDD: major depressive disorder; HC: healthy control.
Table 5
Performance of different methods in SZ/HC classification with 10-fold cross-validation.

<table>
<thead>
<tr>
<th>Method</th>
<th>ACC(%)</th>
<th>SEN(%)</th>
<th>SPE(%)</th>
<th>F1(%)</th>
<th>AUC(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nearest Neighbors</td>
<td>64.6(0.6)</td>
<td>69.1(0.8)</td>
<td>61.6(0.6)</td>
<td>61.0(0.8)</td>
<td>64.8(0.6)</td>
</tr>
<tr>
<td>AdaBoost</td>
<td>72.3(1.1)</td>
<td>72.4(1.1)</td>
<td>72.1(1.1)</td>
<td>72.8(1.0)</td>
<td>72.2(1.1)</td>
</tr>
<tr>
<td>Naive Bayes</td>
<td>69.6(0.5)</td>
<td>67.5(0.4)</td>
<td>72.7(0.6)</td>
<td>72.1(0.5)</td>
<td>69.5(0.5)</td>
</tr>
<tr>
<td>Gaussian Process</td>
<td>64.3(0.8)</td>
<td>68.5(1.0)</td>
<td>61.4(0.7)</td>
<td>60.9(0.9)</td>
<td>64.4(0.8)</td>
</tr>
<tr>
<td>Linear SVM</td>
<td>77.1(0.9)</td>
<td>77.4(1.0)</td>
<td>76.8(0.9)</td>
<td>77.4(0.9)</td>
<td>77.1(0.9)</td>
</tr>
<tr>
<td>Deep Neural Net</td>
<td>80.3(0.5)</td>
<td>80.8(0.9)</td>
<td>79.8(0.9)</td>
<td>80.5(0.5)</td>
<td>80.3(0.5)</td>
</tr>
<tr>
<td>GAN</td>
<td>82.1(0.7)</td>
<td>78.1(1.5)</td>
<td>86.2(1.1)</td>
<td>81.6(0.8)</td>
<td>82.3(0.7)</td>
</tr>
</tbody>
</table>

Table 5 The performance of different methods in SZ/HC classification with 10-folds cross-validation. SVM, support vector machine; No FM, No feature matching; GAN, Generative Adversarial Networks; ACC, Accuracy; SEN, sensitivity; SPE, specificity; F1, F-score; AUC, area under curve; SZ: schizophrenia; HC: healthy control.
Table 6

Performance of the leave-one-site-out methods in SZ/HC classification.

<table>
<thead>
<tr>
<th>Method</th>
<th>ACC(%)</th>
<th>SEN(%)</th>
<th>SPE(%)</th>
<th>f1(%)</th>
<th>AUC(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nearest Neighbors</td>
<td>64.3(5.9)</td>
<td>67.7(8.2)</td>
<td>61.8(9.8)</td>
<td>61.7(6.0)</td>
<td>64.4(5.2)</td>
</tr>
<tr>
<td>AdaBoost</td>
<td>70.4(5.0)</td>
<td>70.0(7.8)</td>
<td>70.8(10.0)</td>
<td>71.4(4.2)</td>
<td>70.3(5.0)</td>
</tr>
<tr>
<td>Naive Bayes</td>
<td>66.6(6.0)</td>
<td>64.4(8.8)</td>
<td>69.9(10.9)</td>
<td>69.8(5.1)</td>
<td>66.4(5.7)</td>
</tr>
<tr>
<td>Gaussian Process</td>
<td>61.8(4.0)</td>
<td>64.6(6.5)</td>
<td>59.7(8.7)</td>
<td>59.2(5.2)</td>
<td>61.9(3.1)</td>
</tr>
<tr>
<td>Linear SVM</td>
<td>74.9(2.9)</td>
<td>74.1(7.9)</td>
<td>75.8(8.6)</td>
<td>75.8(1.9)</td>
<td>74.9(3.1)</td>
</tr>
<tr>
<td>Deep Neural Net</td>
<td>77.4(2.5)</td>
<td>77.0(9.0)</td>
<td>77.8(8.3)</td>
<td>78.0(1.8)</td>
<td>77.3(2.4)</td>
</tr>
<tr>
<td>GAN</td>
<td>80.7(3.8)</td>
<td>81.8(3.7)</td>
<td>79.7(9.7)</td>
<td>80.8(3.9)</td>
<td>80.7(3.4)</td>
</tr>
</tbody>
</table>

Table 6 The performance of the leave-one-site-out method in SZ/HC classification. SVM, support vector machine; GAN, Generative Adversarial Networks; ACC, Accuracy; SEN, sensitivity; SPE, specificity; F1, F-score; AUC, area under curve; SZ: schizophrenia; HC: healthy control.
<table>
<thead>
<tr>
<th>Brain Region</th>
<th>FNC node</th>
<th>Brain Region</th>
<th>FNC node</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUD</td>
<td>inferior temporal gyrus (ITG)</td>
<td>FN</td>
<td>middle frontal gyrus (MFG)</td>
</tr>
<tr>
<td>AUD</td>
<td>middle temporal gyrus (MTG)</td>
<td>FN</td>
<td>superior frontal gyrus (SFG)</td>
</tr>
<tr>
<td>DMN</td>
<td>anterior cingulate (AC)</td>
<td>CC</td>
<td>inferior parietal lobe (IPL)</td>
</tr>
<tr>
<td>DMN</td>
<td>anterior cingulate (AC)</td>
<td>CC</td>
<td>superior temporal gyrus (STG)</td>
</tr>
<tr>
<td>SMN</td>
<td>postcentral gyrus (PoCG)</td>
<td>CC</td>
<td>inferior parietal lobe (IPL)</td>
</tr>
<tr>
<td>DMN</td>
<td>Precuneus</td>
<td>CB</td>
<td>fusiform gyrus (FFG)</td>
</tr>
<tr>
<td>SMN</td>
<td>medial frontal gyrus (MeFG)</td>
<td>CB</td>
<td>declive</td>
</tr>
</tbody>
</table>

Table 7 The most contributing FNC and brain region with the leave-one-FNC-out method for MDD vs HC. For corresponding connections, the brain region and FNC node are presented. AUD, auditory; CC, cognitive control; SMN, sensorimotor network; DMN, default-mode network; FN, frontal network; and CB, cerebellum.
Table 8
The most contributing FNC with the leave-one-FNC-out method for SZ vs HC.

<table>
<thead>
<tr>
<th>Brain Region</th>
<th>FNC node</th>
<th>Brain Region</th>
<th>FNC node</th>
</tr>
</thead>
<tbody>
<tr>
<td>SMN</td>
<td>postcentral gyrus (PoCG)</td>
<td>DMN</td>
<td>precuneus</td>
</tr>
<tr>
<td>VIS</td>
<td>middle occipital gyrus (MOG)</td>
<td>DMN</td>
<td>precuneus</td>
</tr>
<tr>
<td>VIS</td>
<td>middle occipital gyrus (MOG)</td>
<td>DMN</td>
<td>angular gyrus (AG)</td>
</tr>
<tr>
<td>SC</td>
<td>putamen</td>
<td>DMN</td>
<td>precuneus</td>
</tr>
<tr>
<td>SMN</td>
<td>precentral gyrus (PreCG)</td>
<td>DMN</td>
<td>angular gyrus (AG)</td>
</tr>
<tr>
<td>AUD</td>
<td>superior temporal gyrus (STG)</td>
<td>VIS</td>
<td>fusiform gyrus (FFG)</td>
</tr>
</tbody>
</table>

Table 8 The most contributing FNC and brain region with the leave-one-FNC-out method for SZ vs HC. For corresponding connections, the brain region and FNC node are presented. SMN, sensorimotor network; DMN, default-mode network; VIS, visual; SC, sub-cortical; and AUD, auditory.